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## AMENDMENT TO THE CLAIMS

This listing of claims will replace all prior versions, and listings, of claims in the application:

- 1. (Currently amended) An oral pharmaceutical dosage form comprising:
  - (a) a core material <u>comprising</u> [that contains] a proton pump inhibitor, at least one [or more] alkaline reacting <u>compound</u> [compound(s)] and optionally pharmaceutically acceptable excipients [having],
  - (b) a water soluble separating layer, and
  - (c) a [an enterie] coating layer comprising at least one enteric polymer,

wherein, [characterized in that] the core material is alkaline reacting, and upon application of the coating layer on the core material, [that] the separating layer is [being] formed in situ [during the enteric coating] as a water soluble salt product between the enteric polymer [coating layer polymer(s)] and the alkaline reacting compound [compound(s)].

- 2. (Currently amended) The [A] dosage form according to claim 1, wherein the alkaline reacting compound is [compounds are] selected from the group consisting of an alkaline reacting organic compound [substances], a hydroxide [hydroxides] of an alkali metal, an [metals or one of their] alkaline salt [salts] of phosphoric acid, an alkaline salt of carbonic acid, an alkaline salt of [or] silicic acid, and [or] an alkaline ammonium salt.
- 3. (Currently amended) The [A] dosage form according to claim 2, wherein the alkaline reacting compound [substance] is selected from the group consisting of a hydroxide of an alkali metal, [of] an alkaline salt of phosphoric acid, an alkaline salt of carbonic acid, an alkaline salt of [of] silicic acid, and [of] an alkaline ammonium salt.
- 4. (Currently amended) <u>The [A]</u> dosage form according to claim 2, wherein the alkaline reacting [compound is an alkaline] organic compound is [substance, e.g.] an amino acid or a salt thereof [, an alkaline amine or a derivative thereof, or an alkaline salt of a weak organic acid].

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- 5. (Currently amended) The [A] dosage form according to claim 3 [2], wherein the [alkaline organic substance is an] amino acid is selected from the group consisting of [, e.g.] lysine, arginine, ornitine and [or] histidine [, or an alkaline amine or a derivative thereof, e.g. N-methyl-p-glueamine or trometamine].
- 6. (Currently amended) The [A] dosage form according to claim 1, wherein the alkaline reacting compound is [compounds are] present in a concentration of more than 0.1 mmol/g dry ingredients in the alkaline containing part of the core material.
- 7. (Currently amended) <u>The [A] dosage form according to claim 1, wherein the enteric polymer is a [coating polymer(s) is/are]</u> hydroxypropyl cellulose <u>derivative</u> [derivative(s), e.g. hydroxypropylmethylcellulose acetate succinate].
- 8. (Currently amended) <u>The [A]</u> dosage form according to claim 1, wherein the enteric coating polymer is a copolymer of methacrylic acid or methylmethacrylate ester [copolymerized methacrylic acid/methacrylic acid/methyl esters].
- 9. (Currently amended) The [A] dosage form according to claim 1, wherein the proton pump inhibitor is a compound of the general formula I or a pharmaceutically acceptable salt thereof or a pure enantiomer thereof in neutral form or in the form of an alkaline salt

wherein

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Het<sub>1</sub> is

$$R_1$$
 $R_2$ 
 $R_3$ 
or
 $R_6$ 

Het2 is

$$R_6$$
 $R_7$ 
 $R_8$ 
 $R_8$ 
 $R_9$ 
 $R_9$ 

X =

wherein N in the benzimidazole moiety means that one of the carbon atoms substituted by R6-R9 [optionally] may be exchanged for a nitrogen atom without any substituents;

R<sub>1</sub>, R<sub>2</sub> and R<sub>3</sub> are the same or different and selected from the group consisting of hydrogen, alkyl, unsubstituted alkoxy, alkoxy [optionally] substituted by fluorine, alkythio, alkoxyalkoxy, dialkylamino, piperidino, morpholino, halogen, phenyl and phenylalkoxy;

R<sub>4</sub> and R<sub>5</sub> are the same or different and selected from hydrogen, alkyl and aralkyl;

R6' is sclected from the group consisting of hydrogen, halogen, trilluoromethyl, alkyl and alkoxy;

 $R_6$ - $R_9$  are the same or different and selected from the group consisting of hydrogen, alkyl, alkoxy, halogen, halo-alkoxy, alkylcarbonyl, alkoxycarbonyl, oxazolyl, and trifluoroalkyl, or adjacent groups  $R_6$ - $R_9$  form ring structures which may be further substituted;

R<sub>10</sub> is hydrogen or forms an alkylene chain together with R<sub>3</sub> and

R<sub>11</sub> and R<sub>12</sub> are the same or different and selected from the group consisting of hydrogen, halogen, [e+] alkyl and [alkyl groups,] alkoxy, which alkyl or alkoxy [groups and moities thereof] may be branched or a [and] straight C<sub>1</sub>-C<sub>9</sub>-chain [chains] or a [comprise] cyclic alkyl [groups, for example cycloalkylalkyl].

- 10. (Currently amended) The [A] dosage form according to claim 1, wherein the proton pump inhibitor is omegrazole or an alkaline salt thereof.
- 11. (Currently amended) <u>The [A]</u>dosage form according to claim 1, wherein the proton pump inhibitor is a pure enantiomer of omegrazole or an alkaline salt thereof.
- 12. (Currently amended) The [A] dosage form according to claim 1, wherein the proton pump inhibitor is lansoprazole, one of its pure enantiomers or a pharmaceutically acceptable salt thereof.
- 13. (Currently amended) The [A] dosage form according to claim 1, wherein the proton pump inhibitor is pantoprazole, one of its pure enantiomers or a pharmaceutically acceptable salt thereof.
- 14. (Currently amended) The [A] dosage form according to claim 1, wherein the [alkaline reacting] core material is in the form of individual pellets [intended for a capsule formulation or a tableted multiple unit dosage form].
- 15. (Currently amended) The [A] dosage form according to claim 1, wherein the |alkaline reacting| core material is a tablet.

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- 16. (Currently amended) The [A] dosage form according to claim 14 [1], wherein individually enteric coated pellets are compressed into a tableted multiple unit dosage form.
- 17. (Currently amended) A process for the preparation of an oral, enteric coated pharmaceutical dosage form comprising the steps of:

forming a core material comprising [that contains] a proton pump inhibitor, at least one [or more] alkaline reacting compounds and optionally pharmaceutically acceptable excipients, and applying a coating layer comprising at least one enteric polymer so as to surround the core material thereby forming in situ [having a water soluble] separating layer as a water soluble product between the alkaline compound and the enteric polymer [and an enteric coating layer characterized in that an alkaline reacting core material is prepared and coated with an enteric coating polymer wherein a separating layer between the core material and the enteric coating layer is formed in situ by a reaction between the enteric coating polymer(s) and the alkaline reacting compound(s) in the core material during the application of the enteric coating onto the alkaline reacting core material].

- 18. (Canceled)
- 19. (Currently amended) A method for inhibiting gastric acid secretion comprising [in mammals and man by] administering [to a host in need thereof a dosage form comprising] a therapeutically effective amount of a dosage form [dose of a proton pump inhibitor] as defined in any of claims 1-16 to a patient in need thereof.
- 20. (Canceled)
- 21. (New) The dosage form according to claim 2, wherein the alkaline reacting organic compound is an alkaline amine or a derivative thereof.

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- 22. (New) The dosage form according to claim 21, wherein the derivative of the alkaline amine is N-methyl-D-glucamine or trometamine.
- 23. (New) The dosage form according to claim 2, wherein the alkaline reacting organic compound is an alkaline salt of a weak organic acid.
- 24. (New) The dosage form according to claim 7, wherein the hydroxypropyl cellulose derivative is hydroxypropylmethylcellulose acetate succinate.